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## **Mental Health and Morbidity of Caregivers and Co-Residents of individuals with Dementia: A Quasi-Experimental Design**

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**Mental Health and Morbidity of Caregivers and Co-Residents of individuals with  
Dementia: A Quasi-Experimental Design**

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## 1 INTRODUCTION

2 With an ageing population and an increased desire for older people to remain in their own  
3 homes the burden on informal caregivers is increasing (Roth et al., 2015b). Historically,  
4 informal caregiving in the USA has been estimated to lead to a saving of approximately  
5 \$2billion per year compared to institutional costs (Leon and Moyer, 1999), but there are  
6 ongoing concerns about the emotional and physical costs to the caregiver (Capistrant et al.,  
7 2014; Schulz and Beach, 1999; Smith et al., 2014). Despite research suggesting that the  
8 economic costs for caring for the terminally ill at home are actually greater than that in  
9 institutional care (Hollander and Chappell, 2007), the majority of the research points to the  
10 cost savings of informal caregiving often overlooking the fact that providing informal care  
11 affects an individual's labour market position and well-being (Hassink and Van den Berg,  
12 2011; Leon and Moyer, 1999; Van Den Berg et al., 2014).

13 Research simultaneously emphasises increased stress and burden associated with informal  
14 caregiving whilst demonstrating an associated mortality advantage (Capistrant et al., 2014;  
15 O'Reilly et al., 2015a; Ramsay et al., 2013; Roth et al., 2015b; Smith et al., 2014). Earlier  
16 reports of the association between caregiving and increased mortality (Schulz and Beach,  
17 1999) have been superseded by more recent studies demonstrating that, compared to non-  
18 caregivers, caregiving is associated with a lower mortality risk (Brown et al., 2009; O'Reilly  
19 et al., 2015a; Ramsay et al., 2013; Roth et al., 2015b; Smith et al., 2014).

20 However, the majority of studies have considered caregivers as a homogenous group and  
21 few have looked at the variation in mental ill-health or mortality risk of the caregiver  
22 according to the health of the care recipient. It is suggested that some caregiving duties,  
23 especially those related to dementia care, may be particularly deleterious to health. This is

important given that the prevalence of dementia is expected to double in next 20 years with an increasing role being played by family caregivers (Ferri et al., 2006). Providing informal care for an individual with dementia is thought to differ from other care scenarios because of the increased demand placed on caregivers due to patient confusion, aggression, personality change and a poor health trajectory (Papastavrou et al., 2012), and the literature generally describes caregiving for individuals with dementia as associated with significantly increased stress, burden and negative mental health consequences (Amirkhanyan and Wolf, 2003; Arai et al., 2014; Ask et al., 2014; Cuijpers, 2005; Loi et al., 2015). However, many of these studies rely on small sample sizes limited to either spousal caregivers (Ask et al., 2014), or parent-child caregivers (Amirkhanyan and Wolf, 2003), others with arguably inappropriate control groups (Roth et al., 2015b). Comparing dementia caregivers to non-caregivers conflates both the travails of caregiving with the emotional burden of having a family member with dementia.

Many researchers have cited Pearlin et al's stress pathway to explain why major chronic challenges such as caregiving can have such negative mental health impacts; but have failed to recognise that this pathway may be true for all family members of individuals with dementia (Pearlin et al., 2011), so the oft quoted association between caregiving for someone with dementia and mental well-being may be as much a consequence of witnessing the suffering/deterioration of a family member with a chronic, disabling illness or of anticipatory bereavement, as the effects of caregiving per se (Brown et al., 2009). On the other hand there is growing literature demonstrating that caregiving can be associated with positive attributes including positive self-concept, role approval and a sense of purpose that

may moderate the effect of caregiving stress (Brown and Brown, 2014; Cohen et al., 2002; Hill and Turiano, 2014). There is therefore a need to separate the effects of caregiving from those of other non-caregiving co-residents.

This study capitalises on a unique data linkage to develop a quasi-experimental design, identifying both caregiving and non-caregiving co-residents of individuals with dementia symptoms, to determine if caregivers fare worse than their non-caregiving co-residents. Specifically the study asks – when compared to co-resident non-caregivers, does providing informal care to a household co-resident with dementia symptoms place additional risk on the likelihood of poor mental health or mortality.

## **METHODS**

### **Data sources**

The Northern Ireland Mortality Study (NIMS) is a prospective record-linkage study derived from the linkage of census returns to subsequently registered deaths. Details of both the NIMS and the linkage processes are described elsewhere (O'Reilly et al., 2012). For this study all March 2011 Census returns were linked with subsequent associated mortality records from the General Register's Office (GRO), following-up until the end of 2013 (33 months of follow-up). The resulting linked data were anonymised, held in a secure environment by the Northern Ireland Statistics and Research Agency (NISRA) and made available to the research team for the purpose of this study. The use of the NIMS for research was approved by the Office for Research Ethics Committees Northern Ireland (ORECNI).

## **Households containing someone with Dementia Symptoms**

Analysis was limited to those living with someone with dementia symptoms. The Census contains information on household structure as well as individuals allowing for the identification of all co-residents of people with dementia symptoms. The 2011 Census contains a health question asking “Do you have any of the following conditions which have lasted, or are expected to last, at least 12 months?” – listing ten conditions including “Frequent periods of confusion or memory loss”. Positive responses to this question were used as a proxy for cognitive decline, termed “dementia symptoms” throughout the remainder of this paper. To minimise misclassification we limited identification of dementia symptoms to those individuals aged 65 years and over. Single item screening questions for dementia have been shown to be effective in informants (Ayalon, 2011; Hendry et al., 2015), and often one person fills in the Census for the household. Over 1.8 million individuals (1,810,863) were enumerated in the 2011 Census of which 19,845 were identified as having dementia symptoms. This corresponds favourably with the Alzheimer’s Society’s estimate for diagnosed/undiagnosed dementia in the Northern Ireland population – estimated at 19,765 in 2013/14 based on the National Health Service Quality and Outcomes Framework indicator from General Practitioner records (Alzheimer’s Society, 2014) – thus reinforcing the criterion validity of this dementia proxy.

## **Co-resident caregivers and non-caregivers**

Co-residents, living with an individual with dementia symptoms, were divided into two categories; caregivers and non-caregivers (Figure 1). The assessment of caregiving responsibilities was derived from the Census question: “Do you look after, or give any help or support to family members, friends, neighbours or others because of either: long-term

physical or mental ill-health/disability?; problems related to old age?” Respondents were instructed not to include anything they did as part of their paid employment. This method has been utilised in previous literature (O’Reilly et al., 2015a; Ramsay et al., 2013). Caregiving intensity was measured by responses to the number of hours care provided each week: none (non-caregiver); caregiving for 1-19 hours; caregiving for more than 20 or more hours per week. While the Census did not ask any further questions about either the care-recipient or the type of care provided, it was assumed that caregivers living with an individual with dementia symptoms were providing care for their ill co-resident. Households with more than one caregiver were included and it was assumed all household caregivers were caring for the ill co-resident. Sensitivity analysis was carried out limited to households with only one caregiver. Those living alone or in a communal establishment were removed from analysis leaving 8,604 individuals with dementia symptoms living with 10,982 caregiving and non-caregiving co-residents (Figure 1).

(Figure 1 about here)

### **Cohort attributes**

Personal characteristics of co-residents were drawn from the Census: including age, gender and marital status (grouped as never married; married; widowed, separated or divorced). As Northern Ireland is an ethnically homogenous country ethnic diversity was summarised as white/non-white. Other cohort attributes known to be associated with poor mental health were identified from the Census including educational attainment (no formal qualifications,

foundation level, completed compulsory schooling up to age 16 years, completed A-levels at age 18 years, and University degree or higher) and socioeconomic status was assessed using a combination of housing tenure and the capital value of the property. Tenure was derived directly from the Census (owner occupiers, private renting or social renting). Capital value had been derived as part of an exercise by central government in 2005 to determine the level of local tax payable by each household. These data were linked and combined with tenure to produce an eight-fold classification defining tenure/taxable value of property (private renting; social renting; and for owner-occupiers, five groups ranging from less than £75k to over £200k) (see Table 1), with a separate category for owners with unvalued homes. Physical health was determined from the limiting long-term illness question, “Are your day to day activities limited because of a health problem or disability which has lasted, or is expected to last, at least 12 months?” Respondents could identify “Yes, limited a lot”, “Yes, limited a little” or “No”. This was aggregated to a binary Yes/No outcome, and has been identified in previous work as an accurate indicator of physical health (Cohen et al., 1995).

#### **Area characteristics**

Area level deprivation was assigned using the “Income” domain of the Northern Ireland Index of Multiple Deprivation (NIMDM) which includes an assessment of the proportion of individuals in an area on means-tested benefits (NISRA, 2010). Areas were defined as 890 homogenous groups of approximately 1,900 individuals and were ranked from least deprived to most deprived, then split into equal quintiles. An indicator of urban-rural residence was based on the NISRA classification of Settlements (grouped as urban,



intermediate and rural locales representing settlements of >75,000 people, 2500-75000 and < 2500 people respectively) (NISRA, 2005).

### **Mental Health status**

Individuals were identified as suffering from poor mental health by the health question in the Census “Do you have any of the following conditions which have lasted, or are expected to last, at least 12 months?” Those who identified as suffering from “An emotional, psychological or mental health condition (such as depression or schizophrenia)” were defined as having poor mental health. This indicator has face validity based on its demographic distribution which mirrors known associations with mental health, and predictive validity as it is a strong predictor of death by suicide and has been used in previous research as an indicator of mental well-being (Tseliou et al., 2015).

### **Death**

Death information from official death records was linked probabilistically to Census data, identifying those who died in the follow-up period between April 2011 and December 2013. Methods of these linkages processes are described in detail elsewhere (O’Reilly et al., 2008b).

### **Analysis Strategy**

The design allocated co-residents of individuals with dementia symptoms into caregiving or non-caregiving groups, with the non-caregiving co-residents providing a natural comparator for the additive effects of caregiving on top of the effect of living with an individual with dementia symptoms. Descriptive statistics recorded the socio-demographic characteristics and variations in health status and mortality co-residents. Multi-level regression models compared the risks associated with both poor mental health and mortality of co-resident

carers with co-resident non-carers, adjusting for the clustering of individuals within households. The distribution of mental health or mortality risk may not only be explained by characteristics of the individuals but also by unmeasured characteristics of the household. Multi-level modelling adjusts for this clustering of individuals within household and allows for the calculation of a Variance Partition Co-efficient (VPC) to determine the amount of variation attributable to the second level co-variate, in this instance, household variation. Interactions between independent variables were tested for moderation effects based on strong suggestions from the descriptive statistics and the literature.

## RESULTS

The study sample consisted of 10,982 co-residents of individuals with dementia symptoms, 4,928 (44.9%) non-caregivers and 6,054 (55.1%) caregivers. The mean age of individuals with dementia symptoms was 78.1 years and of co-residents, 59.3 years with the majority of the cohort white (97.6%). Caregiving and non-caregiving co-residents varied according to their demographic characteristics (Table 1). Caregivers were better educated, lived in more affluent households and had better physical and mental health compared to non-caregivers. There were differences between light and more intense caregivers. Those providing intense care (>20 hours per week) tended to be older with a higher proportion of females compared to light caregivers (<20 hours per week) and non-caregiving co-residents. Non-caregiving co-residents tended to live in less deprived areas yet report poorer physical and mental health compared to caregivers. A higher proportion of light caregivers were male, younger, never married, had higher educational attainment and house value, and better mental and physical health compared to both intense caregivers and non-caregivers (Table 1).

(Table 1 about here)

Overall, 12% of non-caregiving co-residents reported chronic poor mental health compared to 6% and 8% for those providing respectively light and more intensive caregiving duties. Multi-level models determined the likelihood of poor mental health in co-resident caregivers compared to co-resident non-caregivers adjusting for the clustering of individuals within households. Caregiving co-residents, as a group, were at no greater risk of poor mental health compared to non-caring co-residents (OR=0.93, 95% CI 0.79, 1.10) [Results available on request] and Table 2 shows the results of the models separating light and intense caregivers. In the unadjusted model co-resident caregivers providing light care were 55% less likely to report poor mental health compared to co-resident non-caregivers (OR=0.45, 95% CI 0.32, 0.61), and caregivers providing more intense care 44% less likely to report poor mental health (OR=0.56, 95% CI 0.46, 0.67). Adjusting for age, gender, marital status, ethnicity, education and socio-economic status (Model 3) attenuated the relationship slightly but the protective association between caregiving and mental health remained. After adjusting for baseline physical health (Model 4) the association disappeared and there was no significant difference in likelihood of poor mental health between co-resident caregivers and non-caregivers. The amount of variation attributable to the clustering was calculated using the variance partition co-efficient (VPC). The VPC in the fully adjusted model (Model 5) suggests approximately 32% of the variance in propensity to having poor mental health is attributable to household differences, highlighting a substantial influence of household on mental health.

Given the observed disparity in gender distribution of caregiving roles, possible interactions between gender and care provision were tested using likelihood ratio tests with no significant interaction (LR test=5.44 P=0.07).

(Table 2 about here)

Multi-level models were constructed to determine the risk of mortality in co-resident caregivers compared to co-resident non-caregivers adjusting for the clustering of individuals within households. Overall, caregiving co-residents were 33% less likely to die during the study period than non-caring co-residents after adjustment for all characteristics described in Table 1 (OR=0.67, 95% CI 0.56, 0.81) [Results available on request]. Table 3 shows the results of the models separating light and intense caregivers. In the unadjusted model individuals providing light care (<20 hours/week) are 83% less likely to die compared to those not providing care (OR=0.17, 95% CI 0.06, 0.44). Providing intense care resulted in a 59% reduced likelihood of death compared to co-resident non-caregivers (OR=0.41, 95% CI 0.26, 0.67). After full adjustment for factors known to be associated with mortality risk; age, gender, marital status, ethnicity, educational attainment, socio-economic status, physical health, area level deprivation and urbanicity, individuals providing intense care to their co-residents with dementia were 36% less likely to die than those not providing care (OR=0.64, 95% CI 0.53,0.78). Possible interactions between gender and care provision were tested using likelihood ratio tests with no significant interaction between gender and care provision for risk of death (LR test=2.93 p=0.23). The VPC in this fully adjusted model (Model 5) suggests that over and above the factors for which we have already adjusted, only 0.03% of the variance in propensity for death is attributable to household variation.

(Table 3 about here)

## DISCUSSION

This large, representative study demonstrates that: (i) caregivers to those with dementia symptoms are no more likely than co-resident non-caregivers to report having poor mental health, and (ii) caregiving is associated with a lower mortality risk when compared to non-caregiving co-residents. That caregiving is not associated with any additional risk of poor mental health supports Brown *et al.*'s recent recommendation for caution before concluding that the negative effects observed in caregiving are due to the provision of care to another person (Brown and Brown, 2014) and suggests a need for researchers to be more careful about their choice of comparator in studies of informal caregiving. Our findings concur with an earlier USA study that showed that having a close family member, such as a spouse/elderly parent, with dementia or other serious disability can lead to stress or depressive symptoms regardless of whether or not the unaffected family member is providing care (Amirkhanyan and Wolf, 2003). Furthermore, even in these difficult circumstances, not everyone may be affected, and Vitaliano *et al* in a unique twin-study suggests that the relationship between caregiving and psychological distress may be determined by a vulnerability largely shaped by genetic and early life factors (Vitaliano et al., 2014). Results from the multi-level models in our study purport that 33% of the variation in mental health is due to household factors, which would explain why household stressors such as living with someone with dementia symptoms could have the same influence on both caregiving and non-caregiving household members.

249

250 This study clearly suggests that being a caregiver for someone with dementia is associated  
251 with a lower mortality risk than experienced by other co-residents and although this is at  
252 odds with earlier studies (Schulz and Beach, 1999; Schulz et al., 1995), it is in keeping with  
253 more recent studies reporting a significant mortality advantage associated with caregiving  
254 (Brown et al., 2009; Fredman et al., 2010; O'Reilly et al., 2015b, 2008a; Ramsay et al., 2013;  
255 Roth et al., 2015b, 2013) and with Brown et al. who showed that providing more than 14  
256 hours care per week is associated with decreased mortality risk independent of the care  
257 demands of the care recipient (Brown et al., 2009). The reason for the lower mortality risk  
258 amongst caregivers is not entirely clear but may be related to feelings of usefulness that  
259 may shape health trajectories in older adults (Gruenewald et al., 2007), and non-caring co-  
260 residents do not get the rewards associated with caring such as satisfaction, sense of  
261 purpose or the patient-carer bond. Prosocial behaviour has been hypothesized to be part of  
262 a chain of related biochemical events that function to reduce stress and inflammation and,  
263 thereby, promote health; the caregiving-related release of oxytocin, in interaction with  
264 progesterone, promotes stress and immune system regulation, thereby enhancing the  
265 helper's health and longevity (Brown and Brown, 2015).

266

## 267 **Potential Limitations**

268 The Census data are comprehensive and, by definition, representative of the entire  
269 population, with complete coverage of deaths and detailed information on individual-level  
270 health and socio-economic status for accurate adjustment in the models. While  
271 representativeness and generalisability of findings are always a concern, the socio-

272 demographic, socioeconomic and health characteristics of caregivers in this study  
273 correspond well with other reports (O'Reilly et al., 2015a; Ramsay et al., 2013; Smith et al.,  
274 2014), endorsing the reliability of these results. Northern Ireland's dementia care strategy  
275 aims to help individuals remain in their own home as long as possible and like the majority  
276 of countries in Europe, home care is the predominant care setting (DHSSPS, 2011; Rodrigues  
277 et al., 2012). One major limitation of this study lies in assumptions derived of the data.  
278 While the indicator of dementia symptoms is a self-reported measure of "*experience of*  
279 *confusion or memory loss*" (i.e. not clinically diagnosed dementia), it correlates very well to  
280 independent population estimates (Alzheimer's Society, 2014), has reasonable face validity  
281 and by limiting potential care recipients to those aged 65 years and over we can be more  
282 confident that these symptoms are associated with dementia. We also assumed that  
283 caregivers within a household are providing care for their ill co-residents but note that this  
284 methodology has been used successfully elsewhere (O'Reilly et al., 2015a; Ramsay et al.,  
285 2013).

286 Another limitation is that we cannot be sure that co-residents who did not identify as  
287 caregivers are not undertaking any caring duties, i.e. they may be providing care but do not  
288 see their roles as an "informal caregiver" but just undertaking normal family duties. This  
289 potential misclassification would not however alter our main conclusion that there is a need  
290 to include the entire family in any support given to the family of individuals with dementia,  
291 not merely those who identify as the main caregiver. The healthy caregiver effect is a major  
292 concern and although there is some evidence that the people who undertake a caregiving  
293 role are healthier than those who do not, the relatively small change in odds ratios (Table 3,  
294 Model 4) after adjustment for health status at baseline suggests that the associated

mortality advantage is not primarily due to health selection. Finally, we acknowledge that the cross-sectional nature of some outputs of the study do not allow consideration of the dynamic nature of caring which has fluctuations in demand and burden.

## **Implications**

The implications of this study are twofold. Firstly, the results add weight to current recommendations for the need to reassess how the risks and benefits associated with caregiving are perceived and reported both in the scientific and in the popular press (Roth et al., 2015a). Secondly, it highlights an often overlooked at-risk group, those non-caregiving co-residents of individuals with dementia symptoms. Whilst previous work focussing on the mental health of dementia “patient-carer” dyads has attributed most of the observed adverse consequences to the provision of care, it may be that having a family member with dementia, and not the caregiving *per se*, is the pertinent risk factor for poor mental health.

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317

318     **Conflict of interest:**

319     No Disclosures to Report

320

321     **Ethics:**

322     This study was approved by the Office for Research Ethics Committees Northern Ireland.

**Table 1: Characteristics of dementia patient co-residents (n=10,982)**

		<b>Non- carers (4,928)</b>	<b>Light caring responsibility (1,155)</b>	<b>Heavy caring responsibility (4,899)</b>	<b>All (10,982)</b>
<b>Age</b>	<b>Mean (years)</b>	57.6yrs	51.7yrs	62.7yrs	59.3yrs
<b>Gender (%)</b>	<b>Male</b>	52.7	59.3	43.5	49.3
	<b>Female</b>	47.3	40.7	56.5	50.7
<b>Marital Status (%)</b>	<b>Never married</b>	38.7	44.5	25.3	33.3
	<b>Married/Co-habiting</b>	53.8	48.0	68.1	59.6
	<b>Separated/Divorced/Widowed</b>	7.6	7.5	6.5	7.1
<b>Household Size (%)</b>	<b>2 people</b>	45.2	33.7	58.1	49.8
	<b>3 people</b>	26.6	27.7	23.4	25.3
	<b>4 people</b>	12.0	14.5	8.5	10.7
	<b>5+ people</b>	16.1	24.2	10.0	14.2
<b>Education (%)</b>	<b>No qualifications</b>	55.6	25.2	46.8	48.4
	<b>Foundation</b>	17.1	18.7	18.6	17.9
	<b>Compulsory Schooling</b>	9.0	13.9	9.8	9.9
	<b>A levels</b>	6.7	12.7	6.4	7.2
	<b>Degree+</b>	11.7	29.4	18.4	16.6
<b>House Tenure (%)</b>	<b>Social Renting</b>	17.9	9.0	14.6	15.5
	<b>Private Renting</b>	7.2	5.5	7.0	6.9
	<b>Missing</b>	5.1	8.1	4.8	5.3
	<b>&lt;£75,000</b>	13.4	6.4	11.2	11.7
	<b>£75,000 – £99,999</b>	17.2	16.2	17.4	17.2
	<b>£100K - £149,999</b>	22.2	25.4	22.8	22.8
	<b>£150k - £199,999</b>	10.4	14.5	12.5	11.8
	<b>£200K+</b>	6.7	15.0	9.7	8.9
<b>Illness (%)</b>	<b>LLTI</b>	56.3	29.9	45.4	48.7
<b>Deprivation (%)</b>	<b>Deprived</b>	16.0	21.4	20.5	18.6
	<b>2</b>	16.7	22.1	18.8	18.2
	<b>3</b>	20.9	18.4	18.7	19.6
	<b>4</b>	21.3	18.5	20.7	20.7
	<b>Affluent</b>	25.2	19.6	21.4	22.9
<b>Urban (%)</b>	<b>Rural</b>	31.3	36.2	28.5	30.0
	<b>Intermediate</b>	30.7	28.8	29.8	30.1
	<b>Urban</b>	36.7	31.9	40.3	38.7
	<b>Missing</b>	1.3	3.1	1.4	1.5
<b>Health (%)</b>	<b>Psychological or Emotional problem</b>	12.1	6.4	8.4	9.8
	<b>Died</b>	6.4	3.9	4.2	5.1

**Table 2: Multi-level logistic regression model illustrating the likelihood of poor mental health in dementia patient's co-residents, given informal care-giving status; adjusting for the clustering of individuals within households. Figures represent Odds Ratios & 95% Confidence Intervals**

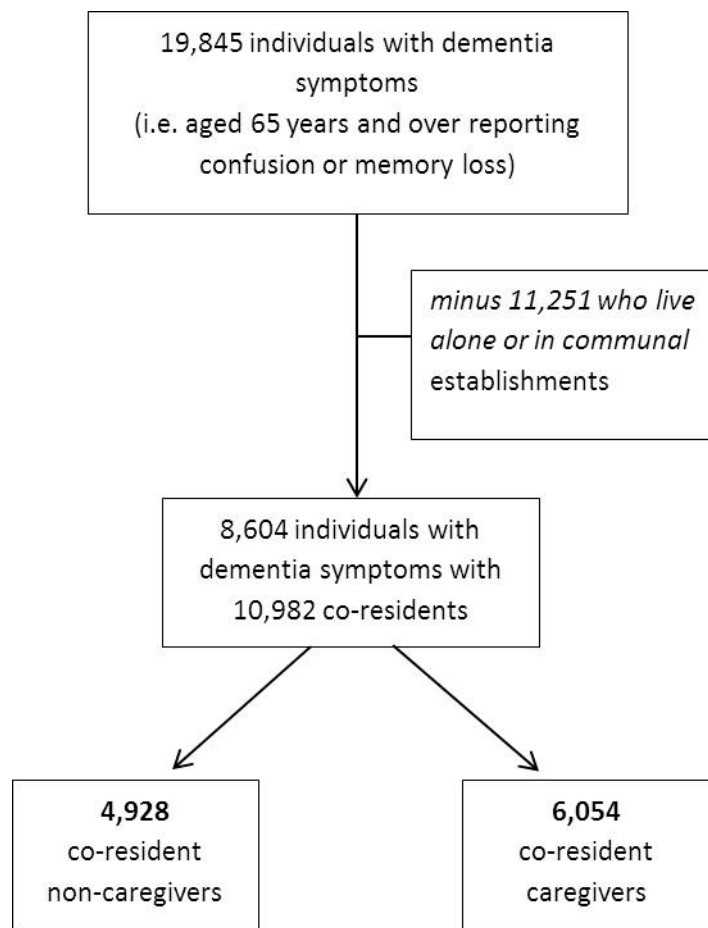
	Co-resident status	Unadjusted	Model 1 Adjusted age & gender	Model 2 +marital status, ethnicity & HH size	Model 3 + SES (tenure & education)	Model 4 + baseline health (LLTI)	Model 5 + area deprivation & urbanicity
	Not a carer	1.00	1.00	1.00	1.00	1.00	1.00
	Light care (<20hrs)	0.45 (0.32,0.61)	0.41 (0.30,0.56)	0.42 (0.31,0.57)	0.52 (0.38,0.71)	0.78 (0.57, 1.07)	0.78 (0.57, 1.08)
	Heavy care (>20 hrs)	0.56 (0.46,0.67)	0.60 (0.50,0.72)	0.59 (0.50,0.71)	0.65 (0.55,0.76)	0.96 (0.81, 1.15)	0.97 (0.81, 1.15)
Gender	Male		1.00	1.00	1.00	1.00	1.00
	Female		1.17 (0.99,1.38)	1.23 (1.05,1.45)	1.27 (1.08,1.49)	1.20 (1.02,1.41)	1.20 (1.02,1.41)
Age	Age (continuous)		0.98 (0.97,0.99)	0.98 (0.97,0.99)	0.98 (0.97,0.99)	0.96 (0.95,0.96)	0.96 (0.95,0.96)
Marital	Never married			1.00	1.00	1.00	1.00
Status	Married/Co-habit			1.30 (1.02,1.66)	1.26 (0.99,1.60)	1.38 (1.07,1.76)	1.37 (1.07,1.76)
	Sep/Div/Wid			2.09 (1.55,2.82)	1.82 (1.35,2.45)	2.00 (1.47,2.74)	1.97 (1.44,2.69)
Ethnicity	White			1.00	1.00	1.00	1.00
	Non-white			0.38 (0.16,0.87)	0.43 (0.19,1.00)	0.68 (0.29,1.56)	0.68 (0.29,1.58)
Household	2 people			1.00	1.00	1.00	1.00
Size	3 people			0.84 (0.68,1.03)	0.96 (0.78,1.18)	1.00 (0.81,1.23)	1.00 (0.81,1.23)
	4 people			0.54 (0.39,0.75)	0.66 (0.48,0.91)	0.75 (0.54,1.03)	0.75 (0.54,1.04)
	5+ people			0.40 (0.29,0.56)	0.54 (0.39,0.75)	0.74 (0.54,1.03)	0.74 (0.54,1.04)
Tenure	Social Renting				1.00	1.00	1.00
	Private Renting				0.54 (0.39,0.77)	0.60 (0.42,0.85)	0.64 (0.45,0.92)
	Missing				0.23 (0.14,0.38)	0.31 (0.18,0.51)	0.32 (0.19,0.56)
	<£75,000				0.34 (0.25,0.47)	0.43 (0.32,0.59)	0.45 (0.33,0.61)
	£75,000 – £99,999				0.47 (0.36,0.61)	0.56 (0.43,0.73)	0.59 (0.45,0.77)
	£100K - £149,999				0.39 (0.30,0.50)	0.47 (0.36,0.61)	0.52 (0.39,0.68)
	£150k - £199,999				0.31 (0.22,0.44)	0.39 (0.28,0.58)	0.45 (0.31,0.64)
	£200K+				0.27 (0.18,0.40)	0.37 (0.25,0.56)	0.43 (0.28,0.67)
Education	No qualifications				1.00	1.00	1.00
	Foundation				1.42 (1.09,1.85)	0.85 (0.64,1.12)	0.84 (0.63,1.10)
	Compulsory School				1.30 (0.97,1.75)	1.07 (0.79,1.45)	1.05 (0.78,1.43)
	A levels				1.23 (0.89,1.72)	1.04 (0.74,1.47)	1.03 (0.73,1.45)

Illness	Degree+				0.97 (0.66,1.41)	0.91 (0.62,1.35)	0.91 (0.61,1.34)
	No					1.00	1.00
	Yes					17.53 (13.29,23.12)	17.51 (13.28,23.10)
Deprivation	Deprived						1.00
	2						0.83 (0.60,1.13)
	3						1.06 (0.78,1.44)
	4						1.12 (0.83,0.51)
Urban	Affluent						1.23 (0.91,1.65)
	Rural						1.00
	Intermediate						1.08 (0.86,1.37)
	Urban						1.03 (0.81,1.30)
	Missing						1.38 (0.62,3.08)
Variance		3.20	3.13	2.39	2.12	1.58	1.57
VPC		0.49	0.49	0.42	0.39	0.32	0.32

**Table 3: Multi-level logistic regression model illustrating the likelihood of mortality in dementia patient's co-residents, given informal care-giving status; adjusting for the clustering of individuals within households. Figures represent Odds Ratios & 95% Confidence Intervals**

		Unadjusted	Model 1 Adjusted age & gender	Model 2 +marital status, ethnicity & HH size	Model 3 + SES (tenure & education)	Model 4 + baseline health (LTTI)	Model 5 + area deprivation & urbanicity
	Co-resident status						
	Not a carer	1.00	1.00	1.00	1.00	1.00	1.00
	Light care (<20hrs)	0.17 (0.06,0.44)	0.71 (0.49,1.02)	0.70 (0.49,1.00)	0.77 (0.54,1.12)	0.82 (0.57,1.19)	0.84 (0.58,1.21)
	Heavy care (>20 hrs)	0.41 (0.26,0.67)	0.59 (0.48,0.71)	0.59 (0.48,0.71)	0.60 (0.50,0.73)	0.65 (0.54,0.80)	0.65 (0.53,0.79)
Gender	Male		1.00	1.00	1.00	1.00	1.00
	Female		0.49 (0.41,0.59)	0.49 (0.41,0.59)	0.50 (0.41,0.60)	0.49 (0.41,0.59)	0.48 (0.40,0.59)
Age	Age (continuous)		1.10 (1.09,1.11)	1.10 (1.09,1.11)	1.10 (1.09,1.11)	1.09 (1.08,1.10)	1.09 (1.08,1.10)
Marital Status	Never married		1.00	1.00	1.00	1.00	1.00
	Married/Co-habit		1.12 (0.82,1.52)	1.13 (0.83,1.53)	1.17 (0.85,1.59)	1.21 (0.89,1.66)	
	Sep/Div/Wid		0.83 (0.53,1.29)	0.77 (0.49,1.21)	0.84 (0.54,1.31)	0.81 (0.52,1.27)	
Ethnicity	White		1.00	1.00	1.00	1.00	1.00
	Non-white		0.54 (0.07,4.01)	0.58 (0.08,4.36)	0.61 (0.08,4.65)	0.57 (0.08,4.32)	
Household Size	2 people		1.00	1.00	1.00	1.00	1.00
	3 people		0.94 (0.74,1.20)	0.97 (0.76,1.23)	0.97 (0.76,1.24)	0.98 (0.77,1.25)	
	4 people		0.81 (0.51,1.27)	0.85 (0.54,1.35)	0.86 (0.55,1.37)	0.91 (0.57,1.44)	
	5+ people		1.20 (0.81,1.79)	1.33 (0.88,2.00)	1.42 (0.94,2.15)	1.47 (0.96,2.23)	
Tenure	Social Renting				1.00	1.00	1.00
	Private Renting				0.85 (0.59,1.23)	0.87 (0.61,1.26)	0.94 (0.65,1.36)
	Missing				0.63 (0.38,1.03)	0.68 (0.41,1.13)	0.78 (0.46,1.35)
	<£75,000				0.77 (0.56,1.08)	0.82 (0.59,1.15)	0.87 (0.63,1.22)
Education	£75,000 – £99,999				0.77 (0.57,1.04)	0.81 (0.60,1.10)	0.85 (0.63,1.16)
	£100K - £149,999				0.60 (0.45,0.81)	0.65 (0.48,0.88)	0.70 (0.51,0.95)
	£150k - £199,999				0.71 (0.50,1.00)	0.79 (0.55,1.10)	0.81 (0.56,1.18)
	£200K+				0.65 (0.43,0.99)	0.73 (0.48,1.12)	0.75 (0.48,1.18)
	No qualifications				1.00	1.00	1.00
	Foundation				1.47 (1.06,2.03)	1.32 (0.95,1.82)	1.38 (0.99,1.93)
	Compulsory School				1.50 (1.03,2.19)	1.41 (0.97,2.06)	1.42 (0.97,2.08)
	A levels				1.23 (0.72,2.12)	1.20 (0.70,2.08)	1.22 (0.71,2.12)

Illness	Degree+				1.74 (0.98,3.11)	1.62 (0.94,2.91)	1.63 (0.91,2.92)
	No					1.00	1.00
Deprivation	Yes					2.75 (2.10,3.59)	2.76 (2.11,3.61)
	Deprived						1.00
	2						0.79 (0.57,1.08)
	3						0.89 (0.66,1.21)
Urban	4						0.98 (0.73,1.32)
	Affluent						0.86 (0.64,1.18)
	Rural						1.00
	Intermediate						1.18 (0.91,1.54)
	Urban						1.34 (1.04,1.73)
	Missing						0.98 (0.40,2.34)
Var		263.66	0.01	<0.001	0.001	0.001	0.001
VPC		0.99	0.003	<0.0003	<0.0003	<0.0003	<0.0003



**Figure 1: Identification of cases (co-resident caregivers) and controls (co-resident non-caregivers)**

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